Appl. No.

: 10/530,904

Filed

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April 8, 2005

REMARKS

Applicants have amended the specification to correct a typographical error in the identification of Formula Ia. Due to a formal mistake at the European Patent Office, a copy of amended pages 27 and 28 was not added to the IPER. In support, Applicants submit a copy of the IPER with Annex as prepared by the EPO as Attachment A. Applicants respectfully request entry of the amendment. No new matter is added herewith.

Conclusion

Should there be any questions concerning this application, the Examiner is invited to contact the undersigned agent at the telephone number appearing below. Please charge any additional fees, including any fees for additional extension of time, or credit overpayment to Deposit Account No. 11-1410.

Respectfully submitted,

KNOBBE, MARTENS, OLSON & BEAR, LLP

Dated

By:

Che Swyden Chereskin, Ph.D.

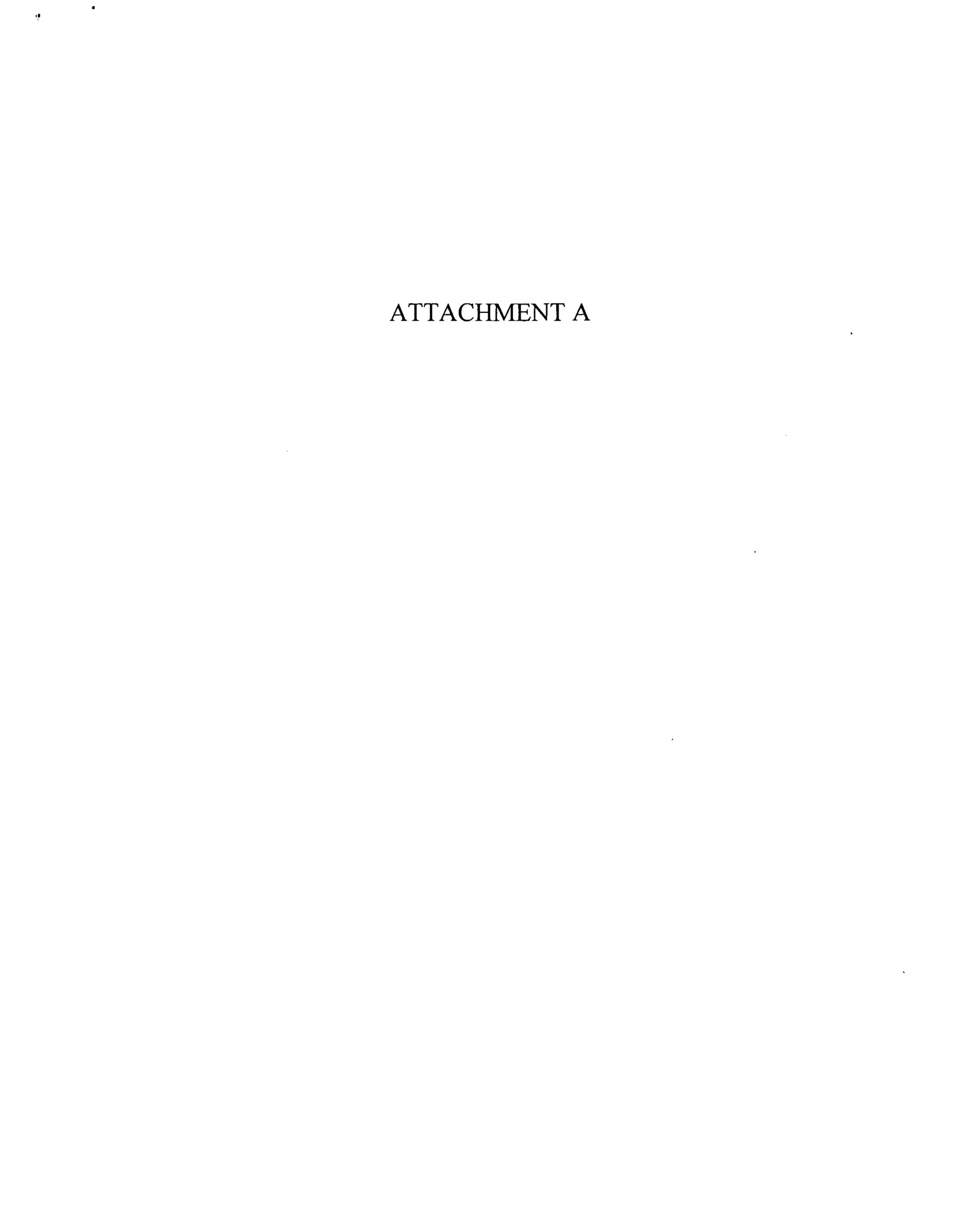
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PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference UNI-001-PCT				FOR FURTHER ACTION See Notification of Transmittal of International Preliminary Examination Report (Form PCT/IPEA/416)			
International application No. PCT/EP 03/11194				International filing date (da 09.10.2003	ay/mor	nth/year)	Priority date (day/month/year) 09.10.2002
1	rnation 7D51		ent Classification (IPC) or bo	oth national classification and	IPC		
	licant IBIOS	SCRE	EN S.A. et al.				
1.	 This international preliminary examination report has been prepared by this International Preliminary Examining Authority and is transmitted to the applicant according to Article 36. 						
2.	2. This REPORT consists of a total of 6 sheets, including this cover sheet.						
	This report is also accompanied by ANNEXES, i.e. sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT).						
	These annexes consist of a total of 19 sheets.						
3.	This	repo	rt contains indications rela	ating to the following item	s:		
	1	\boxtimes	Basis of the opinion				
	11		Priority				
	111	\boxtimes	Non-establishment of o	pinion with regard to nove	elty, ir	nventive step ai	nd industrial applicability
	IV		Lack of unity of inventio	n			
				nder Rule 66.2(a)(ii) with regard to novelty, inventive step or industrial applicability;			
	VI		Certain documents cited	d			
	VII		Certain defects in the in	ternational application			
	VIII		Certain observations on	the international applica	lion		
Date	Date of submission of the demand			D	ato of	completion of this	report
05.0	05.05.2004			2	3.11.	2004	
		exami	g address of the International ining authority:	A	Authorized Officer		
	European Patent Office D-80298 Munich Tel. +49 89 2399 - 0 Tx: 523656			S epmu d	oletti [,]	-Cremers, K	
Fax: +49 89 2399 - 4465				•	elepho	ne No. +49 89 20	399-8541



INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/EP 03/11194

			_
1.	Basis	of the	e report

Description, Pages

1. With regard to the **elements** of the international application (Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17)):

		1,	3-26, 30-73	as originally filed				
		27	, 28	filed with telefax on 30.08.2004				
٠,		2, 2a, 29, 29a		received on 03.11.2004 with letter of 03.11.2004				
)		Claims, Numbers						
		1-3	32	received on 03.11.2004 with letter of 03.11.2004				
		Dra	wings, Sheets					
		1/1	3-13/13	as originally filed				
			With regard to the language, all the elements marked above were available or furnished to this Authority in the language in which the international application was filed, unless otherwise indicated under this item.					
		The	ese elements were av	vailable or furnished to this Authority in the following language: , which is:				
			the language of a translation furnished for the purposes of the international search (under Rule 23.					
	\Box the language of publication of the international application (under Rule 48.3(b)).							
]		the language of a translation furnished for the purposes of international preliminary examination (under Rule 55.2 and/or 55.3).					
,		With regard to any nucleotide and/or amino acid sequence disclosed in the international application, the international preliminary examination was carried out on the basis of the sequence listing:						
	[contained in the inte	ernational application in written form.				
	[filed together with th	ne international application in computer readable form.				
furnished subsequently to this Authority in written form.			furnished subseque	ntly to this Authority in written form.				
	[furnished subseque	ntly to this Authority in computer readable form.				
	[The statement that the subsequently furnished written sequence listing does not go beyond the disclos in the international application as filed has been furnished.						
	[The statement that the listing has been furn	the information recorded in computer readable form is identical to the written sequence ished.				
4	4.	The	amendments have r	esulted in the cancellation of:				
	C		the description,	pages:				
			the claims,	Nos.:				
			the drawings,	sheets:				

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

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	5	. 🗆	This report has been established been considered to go beyor		,	the amendments had not been made, since they have filed (Rule 70.2(c)).		
			(Any replacement sheet contreport.)	taining	such amend	ments must be referred to under item 1 and annexed to this		
	6.	6. Additional observations, if necessary:						
	111	l. No	n-establishment of opinion v	vith re	gard to nov	elty, inventive step and industrial applicability		
	1.		he questions whether the claimed invention appears to be novel, to involve an inventive step (to be non- ovious), or to be industrially applicable have not been examined in respect of:					
)								
		\boxtimes	claims Nos. 32					
			because:					
		\boxtimes	the said international applicat does not require an internation			ms Nos. 32 relate to the following subject matter which mination (specify):		
			see separate sheet					
		L	the description, claims or draw that no meaningful opinion co			icular elements below) or said claims Nos. are so unclear cify):		
			the claims, or said claims Nos. are so inadequately supported by the description that no meaningful opinion could be formed.					
			no international search report	has be	een establish	ed for the said claims Nos.		
2. A meaningful international preliminary examination cannot be carried out due to the failure of amino acid sequence listing to comply with the standard provided for in Annex C of the Adaptive Instructions:				•				
			the written form has not been furnished or does not comply with the Standard.					
			the computer readable form h	as not	been furnish	ed or does not comply with the Standard.		
	٧.	Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement						
	1.	Stat	atement .					
		Nov	elty (N)	Yes: No:	Claims Claims	1-32		
		Inve	entive step (IS)	Yes: No:	Claims Claims	1-32		
		Indu	strial applicability (IA)	Yes: No:	Claims Claims	1-31		
	2.	Citat	tions and explanations					

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see separate sheet

POINT III.

For the assessment of the presently worded claim 32, on the question whether it is industrially applicable, no unified criteria exist in the PCT.

The patentability can also be dependent upon the formulation of the claims. The EPO, for example, does not recognise as industrially applicable claims to the use of a compound in medical treatment, but will allow, however, claims to a known compound for first use in medical treatment and the use of such a compound for the manufacture of a new medical treatment.

POINT V.

The following documents, quoted in the I.S.R., have been considered as relevant for the examination of the present application. Their numbering will be adhered to for the rest of the procedure.

D1: WO-A-98 52562, cited in the application.

D2: J.A.C.S, organic and bio-organic chemistry, 1983, 12, pp. 2827-35.

D3: US-A-5 645 988.

Novelty. 1.

- In view of the fact that the compounds of present invention as claimed are not disclosed in D1 because they possess a saturated spiro-condensed thioazolidine ring instead of the insaturated version of uscharin and that possibly said ring must be substituted, they can be regarded as novel with respect to the content of D1. Moreover, some of the presently claimed compounds differ merely from the uscharin
 - of D1 in that they are merely characterised by the defined substituents R1 as on file and , therefore, they can also be regarded as novel with respect to the content of D1.
- 1.2 Insofar as compounds (1b), (3c) and (3b) of D2 are not part of claimed matter because they are either non substituted derivatives of those (substituted) of the claims, or they

are substituted differently on position 19 (cf. R¹ is different).

Consequently, the claimed matter can be regarded as novel with respect to the content of D2.

1.3 In view of the fact that the compound named 650362 of D3 does not fall within the scope of the claims on file, they can be regarded as novel with respect to its content.

2. Inventiveness.

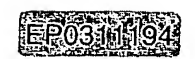
In view of the comparative data which are encompassed in the description and which show the advantages of the claimed compounds in comparison with uscharin, the inventiveness towards D1 and D2 can be acknowledged.

3. Formal Objection.

The attention of the Applicant is already drawn to the fact that he will be faced with an objection towards the content of present claim 23 when the application will reach the European regional proceedings because said claim refers to the description which is not allowable (see Rule 29 (6) EPC) according to the EPC.







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Compositions comprising uscharin or salts thereof have been reported to be usable for treatment of medical conditions related to cell proliferation. For example, US patent No 6,342,490 and WO- 9852562 both describe compositions comprising uscharin or salts thereof and the use of uscharin to combat cell proliferation, e.g. in the treatment of cancer.

Some of the known cardenolide glycosides, f.e. calotropin and uzarigenin, are cytotoxic for cell cultures but are not mentioned to show *in vivo* tumor-inhibiting activity. Also uscharin has been shown to have some cytotoxic activity on tumor cells *in vitro*. In addition, uscharin was also described to have *in vivo* tumor-inhibiting effects, as for instance described in US patent No 6,342,490. Derivatives of uscharin have not been reported so far to be useful for medical applications.

15 Cheung et al. (1983; J. Chem. Soc. Perkin Transactions 1: Organic and bio-organic chemistry (1971-1999) (12) 2827-235) disclose the stereochemistry of cardenolide glycosides of Asclepiadaceae including 19-deoxyuscharin, uscharin and voruscharin.

In US 5,645,988 methods of identifying drugs with selective effects against cancer cells are presented. The drug indicated with 650362 shows some similarity with uscharin.

It is a general object of the present invention to provide novel cardenolide glycosides, which have a cytotoxic activity. It is another general object of the present invention to provide novel cardenolide glycosides, which can be exploited in medical applications.

SUMMARY

In a first aspect, the present invention relates to a compound of the formula I or a pharmaceutically acceptable salt thereof,

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formula !

$$R_4$$
 R_5
 R_2
 R_1
 R_3
 R_4
 R_3

wherein R¹ is selected from the group comprising hydrogen, alkyl, alkenyl, alkynyl, alkyloxy, alkyloxyalkyl, alkylthioalkyl, alkyloxycarbonyl, alkylthiocarbonyl, alkylthiocarbonyl, cycloalkylalkyl, cycloalkylalkoxythiocarbonyl, cycloalkylalkoxythioalkyl, cycloalkylalkoxythioalkyl,

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from the group indicated in above; wherein R² and R³ are hydroxyl and wherein R⁴ and R⁵ are hydrogen or alkyl.

In another preferred embodiment, the invention relates to an uscharin derivative having the formula la, wherein R¹ is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxyalkyl, cycloalkylalkyl, silyloxyalkyl, aralkyl, arylalkenyl, carboxyl, Het¹oxyalkyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het²oxyalkyl, Het²alkyloxyalkyl, Het²aryloxyalkyl, optionally substituted by one or more substituents independently selected from the group indicated above; wherein R² and R³ are hydroxyl and wherein R⁴ and R⁵ are hydrogen.

Another further embodiment of the invention relates to a compound of formula lb, formula lb

$$R_4$$
 R_5
 R_2
 R_1
 R_3
 R_4
 R_4
 R_5
 R_4
 R_4
 R_4
 R_4
 R_5

wherein R1 is selected from the group comprising alkenyl, alkynyl, alkyloxyalkyl, alkylthioalkyl, alkyloxycarbonyl, alkanoyl, cycloalkylalkyl, cycloalkylcarbonyl, cycloalkylalkanoyl, cycloalkylalkoxycarbonyl, cycloalkylthioalkyl, alkylcarbonyloxyalkyl, arylcarbonyloxyalkyl, cycloalkylcarbonyloxyalkyl, silyloxyalkyl, aralkyl, arylalkenyl, arylcarbonyl, aryloxycarbonyl, aralkoxycarbonyl, aryithioalkyl, aralkanoyl, aroyl, silyloxyalkyl, carboxyl, alkenylcarbonyl, Het¹oxyalkyl, Het¹alkoxycarbonyl, Het¹oxycarbonyl, Het¹aryloxyalkyl, alkynylcarbonyl, Het¹aralkoxycarbonyl, Het¹alkyloxyalkyl, Het¹arylthioalkyl, Het¹aryloxycarbonyl, Het¹alkyloxyalkylcarbonyl, Het¹aryloxyalkylcarbonyl, Het¹oxyalkylcarbonyl, Het¹alkylcarbonyloxyalkyl, Het¹aralkylcarbonyloxyalkyl, Het²oxyalkyl, Het¹carbonyloxyalkyl, Het²alkoxycarbonyl, Het²aralkoxycarbonyl, Het²alkyloxyalkyl, Het²oxycarbonyl, Het²oxyalkylcarbonyl, Het²aryloxycarbonyl, Het²aryloxyalkyl, Het²arylthioalkyl, Het²aryloxyalkylcarbonyl, Het²carbonyloxyalkyl, Het²alkyloxyalkylcarbonyl, Het²alkylcarbonyloxyalkyl, Het²aralkylcarbonyloxyalkyl, CR⁸=NR⁷, CR⁶=N(OR⁷),





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29a

with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹ Het¹ alkyl, Het¹ aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R¹ is optionally substituted by one or more substituents independently selected from the group as indicated above,

wherein R^2 and R^3 are hydroxyl and wherein R^4 is replaced by a double bond between the N atom and the C carbon atom of the N-containing heterocyclic ring of formula I; and wherein R^5 is hydrogen.

According to this embodiment, this compound may also be represented by the formula III:





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CLAIMS

 A compound of the formula I or a pharmaceutically acceptable salt thereof, formula I

$$R_4$$
 R_5
 R_2
 R_1
 R_3
 R_3

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wherein R1 is selected from the group comprising hydrogen, alkyl, alkenyl, alkynyl, alkyloxycarbonyl, alkylthiocarbonyl, alkylthioalkyl, alkyloxyalkyl, alkyloxy, cycloalkylthiocarbonyl, cycloalkylaikanoyl, cycloalkylcarbonyl, cycloalkylalkyl, cycloalkylalkoxythiocarbonyl, cycloalkylthioalkyl, cycloalkylalkoxycarbonyl, alkylcarbonyloxyalkyl, arylcarbonyloxyalkyl, cycloalkylcarbonyloxyalkyl, silyloxyalkyl, aralkyl, aralkoxycarbonyl, arylthiocarbonyl, aryloxycarbonyl. arylcarbonyl. arylalkenyl, arylalkylthiocarbonyl, aryloxyalkyl, arylthioalkyl, haloalkyl, hydroxyalkyl, aralkanoyl, aroyl, aryloxycarbonylalkyl, aryloxyalkanoyl, carboxyl, formyl, alkenylcarbonyl, alkynylcarbonyl, Het¹, Het¹aryl, Het¹carbonyl, Het¹aralkyl, Het¹cycloalkyl, Het¹oxyalkyl, Het¹alkyl, Het¹alkoxycarbonyl, Het¹alkylthiocarbonyl, Het¹oxycarbonyl, Het¹thiocarbonyl, Het¹alkanoyl, Het'aralkanoyl, Het'aryloxyalkyl, Het'alkyloxyalkyl, Het'arylthloalkyl, Het'aryloxycarbonyl, Het¹aroyl, Het¹oxyalkylcarbonyl, Het¹alkyloxyalkylcarbonyl, Het¹aralkoxycarbonyl, Het¹carbonyloxyalkyl, Het¹alkylcarbonyloxyalkyl, Het¹aryloxyalkylcarbonyl, Het²aralkyl, Het²alkyloxyalkyl, Het²alkyl; Het²oxyalkyl, Het¹aralkylcarbonyloxyalkyl, Het²carbonyl, Het²oxycarbonyl, Het²thiocarbonyl, Het²alkanoyl, Het²alkylthiocarbonyl, Het²alkoxycarbonyl, Het²aralkanoyl, Het²aralkoxycarbonyl, Het²aryloxycarbonyl, Het²aroyl, Het²alkyloxyalkylcarbonyl, Het²arylthioalkyl, Het²oxyalkylcarbonyl, Het²aryloxyalkyl, Het²alkylcarbonyloxyalkyl, Het²carbonyloxyalkyl, Het²aryloxyalkylcarbonyl, Het²aralkylcarbonyloxyalkyl, cyano, aminocarbonyl, aminoalkanoyl, aminoalkyl, CR⁶=NR⁷ or CR⁶=N(OR⁷), with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹, Het¹alkyl, Het¹aryl, alkenyl, alkynyl, aminoalkyl, arylcarbonylamino, alkylthiocarbonylamino aminoaryl, alkylcarbonylamino, and arylthiocarbonylamino;





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wherein R² and R³ are independently selected from the group comprising hydroxyl, alkyloxy, alkyloxy, arylsilyloxy, alkyloxyalkyloxy, cycloalkyloxy cycloalkylalkyloxy, aralkyloxy, aryloxyalkyloxy, silyloxy, alkylcarbonyloxy, aryloxycarbonyloxy, cycloalkylcarbonyloxy, haloalkyloxy, hydroxyalkyloxy, aralkanoyloxy, arolloxy, aryloxycarbonylalkyloxy, formyloxy, Het¹alkyloxy, Het¹oxyalkyloxy, Het¹aryloxy, Het¹aralkyloxy, Het¹cycloalkyloxy, Het¹carbonyloxy, Het¹aralkanoyloxy, Het¹aralkanoyloxy, Het¹aryloxyalkyloxy, Het²aralkanoyloxy, Het²aralkyloxy, Het²aralkyloxy, Het²aralkyloxy, Het²aryloxyalkyloxy, Het²aryloxy, Het²aryloxyalkyloxy, Het²aryloxy, Het²aryloxy, Het²aryloxy, Het²aryloxy, Het²aryloxy, Het²aryloxy, Het²aryloxy, Het²aryloxy, Het²aryloxy, Het²aryloxy,

wherein R1 R2 and R3 are optionally substituted by one or more substituents independently selected from the group comprising alkyl, aralkyl, aryl, Het1, Het2, cycloalkyl, alkyloxycarbonyl, carboxyl, aminocarbonyl, mono- or di(alkyl)aminocarbonyl, aminosulfonyl, alkylS(=O), hydroxy, cyano, halogen or amino optionally mono- or disubstituted wherein the substituents are independently selected from the group comprising alkyl, aryl, aralkyl, aryloxy, arylamino, arylthio, aryloxyalkyl, arylaminoalkyl, aralkoxy, alkylthio, alkoxy, aryloxyalkoxy, arylaminoalkoxy, aralkylamino, aryloxyalkylamino, arylaminoalkylamino, arylthioalkoxy, aryloxyalkylthio, aralkylthio, arylaminoalkylthio, arylthioalkylamino, arylthioalkylthio, alkylamino, cycloalkyl, cycloalkylalkyl, Het¹, Het², Het¹alkyl, Het²alkyl, Het¹amino, Het²amino, Het¹alkylamino, Het²alkylamino, Het¹thio, Het²thio, Het¹alkylthio, Het²alkylthio, Het¹oxy and Het²oxy, OR⁸, SR⁸, SO₂NR⁸R⁹, SO₂N(OH)R⁸, CN, CR⁸=NR⁹, S(O)R⁸, SO₂R⁸, CR⁸=N(OR⁹), N₃, NO₂, NR⁸R⁹, N(OH)R⁸, C(O)R⁸, C(S)R⁸, CO₂R⁸, C(O)SR⁸, C(O)NR⁸R⁹, C(S)NR⁸R⁹, C(O)N(OH)R9, C(S)N(OH)R8, NR8C(O)R9, NR8C(S)R9, N(OH)C(O)R9, N(OH)C(S)R8, NR⁸CO₂R⁹, NR⁸C(O)NR⁹R¹⁰, and NR⁸C(S)NR⁹R¹⁰, N(OH)CO₂R⁸, NR⁸C(O)SR⁹, N(OH)C(O)NR⁸R⁹, N(OH)C(S)NR⁸R⁹, NR⁸C(O)N(OH)R⁹, NR⁸C(S)N(OH)R⁹, NR⁸SO₂R⁹, NHSO₂NR⁸R⁹, NR⁸SO₂NHR⁹, P(O)(OR⁸)(OR⁹),

with t being an integer between 1 and 2, and R^o R^o and R^{lo} being each independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het^l Alkyl, Het^l aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R⁴ is selected from the group comprising oxo, hydroxyl, alkyl, alkenyl, alkynyl, alkanediyl, alkyloxy, alklylthio, alkylamino, alkyloxyalkyl, arylcarbonylalkyl, alkylcarbonylalkyl, alkanoyl, cycloalkylcarbonylalkyl,





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cycloalkyl, cycloalkyloxy, cycloalkylthio, cycloalkylamino, cycloalkylalkyl, cycloalkylalkanoyl, aralkyl, arylalkenyl, arylcarbonyloxy, aryloxycarbonyloxy, aralkoxycarbonyloxy, haloalkylthio, haloalkylamino , hydroxyalkyl, aralkanoyl, aryloxyalkyl, haloalkyloxy, aryloxycarbonylalkyl, aryloxyalkanoyl, Het¹, Het¹alkyl, Het¹oxy, Het¹oxyalkyl, Het¹aryl, Het¹aralkyl, Het¹cycloalkyl, Het¹aryloxyalkyl, Het¹aroyl, Het², Het²oxy, Het²alkyl; Het²oxyalkyl, Het²aralkyl, Het²cycloalkyl, Het²aryl, Het²alkanoyl, Het²aralkanoyl, Het²aroyl, Het²aryloxyalkyl, aminocarbonyl, aminoalkanoyl, aminoalkyl, optionally substituted by one or more substituents independently selected from the group comprising alkyl, aralkyl, aryl, Het1, Het2, cycloalkyl, alkyloxycarbonyl, carboxyl, aminocarbonyl, mono- or di(alkyl)aminocarbonyl, aminosulfonyl, alkylS(=O), hydroxy, cyano, halogen or amino optionally mono- or disubstituted wherein the substituents are independently selected from the group comprising alkyl, aryl, aralkyl, aryloxy, arylamino, arylthio, aryloxyalkyl, arylaminoalkyl, aralkoxy, alkylthio, alkoxy, aryloxyalkoxy, aylaminoalkoxy, aralkylamino, aryloxyalkylamino, arylaminoalkylamino, arylthioalkoxy, arylaminoalkylthio, arylthioalkylthio, aryloxyalkylthio, aralkylthio, arylthioalkylamino, alkylamino, cycloalkyl, cycloalkylalkyl, Het1, Het2, Het1alkyl, Het2alkyl, Het1amino, Het2amino, Het¹alkylamino, Het²alkylamino, Het¹thio, Het²thio, Het¹alkylthio, Het²alkylthio, Het¹oxy and Het²oxy, OR¹¹, SR¹¹, SO₂NR¹¹R¹², SO₂N(OH)R¹¹, CN, CR¹¹=NR¹², S(O)R¹¹, SO₂R¹¹, $CR^{11}=N(OR^{12})$, N_3 , NO_2 , $NR^{11}R^{12}$, $N(OH)R^{11}$, $C(O)R^{11}$, $C(S)R^{11}$, CO_2R^{11} , $C(O)SR^{11}$, C(O)NR¹¹R¹², C(S)NR¹¹R¹², C(O)N(OH)R¹², C(S)N(OH)R¹¹, NR¹¹C(O)R¹², NR¹¹C(S)R¹², N(OH)C(O)R¹², N(OH)C(S)R¹¹, NR¹¹CO₂R¹², NR¹¹C(O)NR¹²R¹³, and NR¹¹C(S)NR¹²R¹³, N(OH)CO₂R¹¹, NR¹¹C(O)SR¹², N(OH)C(O)NR¹¹R¹², N(OH)C(S)NR¹¹R¹², NR¹¹C(O)N(OH)R¹², $NR^{11}C(S)N(OH)R^{12}$, $NR^{11}SO_2R^{12}$, $NHSO_2NR^{11}R^{12}$, $NR^{11}SO_2NHR^{12}$, $P(O)(OR^{11})(OR^{12})$, wherein t is an integer between 1 and 2, R11, R12 and R13 are each independently selected from the group comprising hydrogen, alkyl, alkenyl, and alkynyl; and

wherein R⁵ is selected from the group comprising hydrogen, oxo, hydroxyl, alkyl, alkenyl, alkynyl, alkanediyl, alkyloxy, alkyloxyalkyl, arylcarbonylalkyl, alkylcarbonylalkyl, alkylcarbonylalkyl, alkylcarbonylalkyl, arylalkenyl, cycloalkylalkyl, cycloalkylalkyl, cycloalkylalkanoyl, aryl, aralkyl, arylalkenyl, arylcarbonyloxy, aryloxycarbonyloxy, aralkoxycarbonyloxy, aryloxyalkyl, haloalkyl, hydroxyalkyl, aralkanoyl, aryloxycarbonylalkyl, aryloxyalkanoyl, Het¹alkyl, Het¹oxy, Het¹aryloxyalkyl, Het¹aryl, Het¹aralkyl, Het¹cycloalkyl, Het¹aryloxyalkyl, Het¹aroyl, Het²aralkyl, Het²aralkyl, Het²cycloalkyl, Het²aryl, Het²aralkanoyl, Het²aralkanoyl, Het²aroyl, Het²aryloxyalkyl, aminocarbonyl, aminoalkanoyl, aminoalkyl, optionally substituted by one or more substituents independently selected from the group comprising alkyl, aralkyl,



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aryl, Het1, Het2, cycloalkyl, alkyloxycarbonyl, carboxyl, aminocarbonyl, mono- or di(alkyl)aminocarbonyl, aminosulfonyl, alkylS(=O), hydroxy, cyano, halogen or amino optionally mono- or disubstituted wherein the substituents are independently selected from the group comprising alkyl, aryl, aralkyl, aryloxy, arylamino, arylthio, aryloxyalkyl, arylaminoalkyl, aralkoxy, alkylthio, alkoxy, aryloxyalkoxy, aylaminoalkoxy, aralkylamino, aryloxyalkylamino, arylaminoalkylamino, arylthioalkoxy, arylthioalkylamino, aralkylthio, aryloxyalkylthio, arylaminoalkylthio, arylthioalkylthio, alkylamino, cycloalkyl, cycloalkylalkyl, Het¹, Het², Het¹alkyl, Het²alkyl, Het¹amino, Het²amino, Het¹alkylamino, Het²alkylamino, Het¹thio, Het²thio, Het¹alkylthio, Het²alkylthio, Het¹oxy and Het²oxy, OR¹¹, SR¹¹, SO₂NR¹¹R¹², SO₂N(OH)R¹¹, CN, CR¹¹=NR¹², S(O)R¹¹, SO₂R¹¹, CR¹¹=N(OR¹²), N₃, NO₂, NR¹¹R¹², N(OH)R¹¹, C(O)R¹¹, C(S)R¹¹, CO₂R¹¹, C(O)SR¹¹, C(O)NR¹¹R¹², C(S)NR¹¹R¹², C(O)N(OH)R¹². C(S)N(OH)R¹¹, NR¹¹C(O)R¹², NR¹¹C(S)R¹², N(OH)C(O)R¹², N(OH)C(S)R¹¹, NR¹¹CO₂R¹², NR¹¹C(O)NR¹²R¹³, and NR¹¹C(S)NR¹²R¹³, N(OH)CO₂R¹¹, NR¹¹C(O)SR¹², N(OH)C(O)NR¹¹R¹². N(OH)C(S)NR¹¹R¹², NR¹¹C(O)N(OH)R¹², NR¹¹C(S)N(OH)R¹², NR¹¹SO₂R¹², NHSO₂NR¹¹R¹², NR¹¹SO₂NHR¹², P(O)(OR¹¹)(OR¹²), wherein t is an integer between 1 and 2, R¹¹, R¹² and R¹³ are each independently selected from the group comprising hydrogen, alkyl, alkenyl, and alkynyl.

2. A compound according to claim 1, having the formula I or a pharmaceutically acceptable salt thereof,

formula I

$$R_4$$
 R_5
 R_2
 R_1
 R_3
 R_4
 R_4
 R_5
 R_5
 R_1
 R_3

wherein R¹ is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxy, alkyloxyalkyl, alkylthioalkyl, alkyloxycarbonyl, alkylthiocarbonyl, alkanoyl, cycloalkylalkyl, cycloalkylalkoxycarbonyl, cycloalkylalkoxythiocarbonyl, cycloalkylthioalkyl, alkylcarbonyloxyalkyl, arylcarbonyloxyalkyl, cycloalkylcarbonyloxyalkyl, arylcarbonyl, aryloxycarbonyl, arylthiocarbonyl, aryloxycarbonyl, arylthioalkyl, arylthioalkyl, arylthioalkyl, haloalkyl, arylthioalkyl, arylthioalkyl, haloalkyl,



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hydroxyalkyl, aralkanoyl, aroyl, aryloxycarbonylalkyl, aryloxyalkanoyl, carboxyl, formyl, alkenylcarbonyl, alkynylcarbonyl, Het1, Het1alkyl, Het1oxyalkyl, Het1aryl, Het1aralkyl, Het¹cycloalkyl, Het¹carbonyl, Het¹alkoxycarbonyl, Het¹alkylthiocarbonyl, Het¹oxycarbonyl, Het¹alkanoyl, Het¹aralkanoyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het¹thiocarbonyl, Het¹arylthioalkyl, Het¹aryloxycarbonyl, Het¹aralkoxycarbonyl, Het¹aroyl, Het¹oxyalkylcarbonyl, Het¹aryloxyalkylcarbonyl, Het¹carbonyloxyalkyl, Het¹alkyloxyalkylcarbonyl, Het²alkyl; Het2oxyalkyl, Het¹aralkylcarbonyloxyalkyl, Het¹alkylcarbonyloxyalkyl, Het²alkyloxyalkyl, Het²aralkyl, Het²carbonyl, Het²oxycarbonyl, Het²thiocarbonyl, Het²alkanoyl, Het²alkoxycarbonyl, Het²aralkanoyl, Het²aralkoxycarbonyl, Het²alkylthiocarbonyl, Het²aryloxycarbonyl, Het²aroyl, Het²aryloxyalkyl, Het²arylthioalkyl, Het²oxyalkylcarbonyl, Het²carbonyloxyalkyl, Het²aryloxyalkylcarbonyl, Het²alkyloxyalkylcarbonyl, Het²alkylcarbonyloxyalkyl, Het²aralkylcarbonyloxyalkyl, cyano, aminocarbonyl, aminoalkanoyl, aminoalkyl, CR6=NR7 or CR6=N(OR7), with R6 and R7 being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het1, Het1alkyl, Het1aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R² and R³ are independently selected from the group comprising hydroxyl, alkyloxy, alkyloxy, arylsilyloxy, alkyloxyalkyloxy, cycloalkyloxy cycloalkyloxy, aralkyloxy, aryloxyalkyloxy, silyloxy, alkylcarbonyloxy, aryloxycarbonyloxy, cycloalkylcarbonyloxy, haloalkyloxy, hydroxyalkyloxy, aralkanoyloxy, aroyloxy, aryloxycarbonylalkyloxy, formyloxy, Het¹alkyloxy, Het¹oxy, Het¹aryloxy, Het¹aryloxy, Het¹aralkyloxy, Het¹cycloalkyloxy, Het¹cycloalkyloxy, Het¹aralkanoyloxy, Het¹aryloxyalkyloxy, Het¹aroyl, Het²oxy, Het²alkyloxy; Het²oxyalkyloxy, Het²aralkyloxy, Het²aralkyloxy, Het²aryloxyalkyloxy, Het²aryloxy, Het²aryloxy,

wherein R¹ R² and R³ are optionally substituted by one or more substituents independently selected from the group comprising alkyl, aralkyl, aryl, Het¹, Het², cycloalkyl, alkyloxycarbonyl, carboxyl, aminocarbonyl, mono- or di(alkyl)aminocarbonyl, aminosulfonyl, alkylS(=O)t, hydroxy, cyano, halogen or amino optionally mono- or disubstituted wherein the substituents are independently selected from the group comprising alkyl, aryl, aralkyl, aryloxy, arylamino, arylthio, aryloxyalkyl, arylaminoalkyl, aralkoxy, alkylthio, alkoxy, aryloxyalkoxy, arylaminoalkoxy, aralkylamino, aryloxyalkylamino, arylaminoalkylamino, arylthioalkylthio, arylthioalkylthio, arylthioalkylthio, arylthioalkylthio, arylthioalkylthio, arylthioalkylthio, arylthioalkylthio,









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alkylamino, cycloalkyl, cycloalkylalkyl, Het¹, Het², Het¹alkyl, Het²alkyl, Het¹amino, Het²amino, Het¹alkylamino, Het²alkylamino, Het¹thio, Het²thio, Het¹alkylthio, Het²alkylthio, Het¹oxy and Het²oxy, OR³, SR³, SO₂NR³R³, SO₂N(OH)R³, CN, CR³=NR³, S(O)R³, SO₂R³, CR³=N(OR³), N₃, NO₂, NR³R³, N(OH)R³, C(O)R³, C(S)R³, CO₂R³, C(O)SR³, C(O)NR³R³, C(S)NR³R³, C(O)N(OH)R³, NR³C(O)R³, NR³C(S)R³, N(OH)C(O)R³, N(OH)C(S)R³, NR³C(O)NR³R¹, NR³C(O)NR³R¹, N(OH)CO₂R³, NR³C(O)SR³, N(OH)C(O)NR³R³, N(OH)C(S)NR³R³, NR³C(O)N(OH)R³, NR³C(S)N(OH)R³, NR³SO₂R³, NHSO₂NR³R³, NR³SO₂NHR³, P(O)(OR³)(OR³),

with t being an integer between 1 and 2, and R⁸ R⁹ and R¹⁰ being each independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹ Alkyl, Het¹ aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R⁴ is oxo and R⁵ is hydrogen or alkyl.

15 3. A compound according to claim 1,

wherein R1 is selected from the group comprising hydrogen, alkyl, hydroxyalkyl, alkenyl, alkynyl, alkyloxyalkyl, alkylthioalkyl, alkyloxycarbonyl, alkanoyl, cycloalkylalkyl, cycloalkylcarbonyl, cycloalkylalkoxycarbonyl, cycloalkylalkanoyl, cycloalkylthioalkyl, alkylcarbonyloxyalkyl, arylcarbonyloxyalkyl, cycloalkylcarbonyloxyalkyl, silyloxyalkyl, aralkyl, arylalkenyl, arylcarbonyl, aryloxycarbonyl, aralkoxycarbonyl, arylthioalkyl, aralkanoyl, aroyl, carboxyl, formyl, alkenylcarbonyl, alkynylcarbonyl, Het¹oxyalkyl, Het¹alkoxycarbonyl, Het¹oxycarbonyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het¹arylthioalkyl, Het¹aryloxycarbonyl, Het¹oxyalkylcarbonyl, Het¹aralkoxycarbonyl, Het¹alkyloxyalkylcarbonyl, Het¹aryloxyalkylcarbonyl, Het¹carbonyloxyalkyl, Het¹alkylcarbonyloxyalkyl, Het¹aralkylcarbonyloxyalkyl, Het²alkyloxyalkyl, Het²oxyalkyl, Het²oxycarbonyl, Het²aralkoxycarbonyl, Het²aryloxycarbonyl, Het²alkoxycarbonyl, Het²aryloxyalkyl, Het²arylthioalkyl, Het²oxyalkylcarbonyl, Het²alkyloxyalkylcarbonyl, Het²aryloxyalkylcarbonyl, Het²carbonyloxyalkyl, Het²alkylcarbonyloxyalkyl, Het²aralkylcarbonyloxyalkyl, CR⁶=NR⁷, $CR^6=N(OR^7)$,

with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹, Het¹alkyl, Het¹aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;



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wherein R² and R³ are independently selected from the group comprising hydroxyl, alkyloxy, alkyloxyalkyloxy, cycloalkyloxy cycloalkylalkyloxy, aralkyloxy, aryloxyalkyloxy, silyloxy, alkylcarbonyloxy, aryloxycarbonyloxy, cycloalkylcarbonyloxy, aryloxycarbonylalkyloxy, formyloxy, Het¹alkyloxy, Het¹oxy, Het¹oxyalkyloxy, Het¹aryloxy, Het¹aralkyloxy, Het¹aralkyloxy, Het¹aralkyloxy, Het¹aralkyloxy, Het¹aralkyloxy, Het²aralkyloxy, Het²aralkyloxy,

wherein R¹ R² and R³ are optionally substituted by one or more substituents independently selected from the group indicated in claim 1; and

wherein R⁴ is selected from the group comprising, oxo, hydroxyalkyl, alkyl, alkenyl, alkylcarbonylalkyl, arylcarbonylalkyl and R⁵ is hydrogen, oxo, hydroxyl, hydroxyalkyl, alkyl, alkyl, alkyl, alkylcarbonylalkyl, arylcarbonylalkyl.

4. A compound according to claim 1 or 2,

wherein R1 is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxyalkyl, alkylthioalkyl, alkyloxycarbonyl, alkanoyl, cycloalkylalkyl, cycloalkylcarbonyl, cycloalkylalkanoyl, cycloalkylalkoxycarbonyl, cycloalkylthioalkyl, alkylcarbonyloxyalkyl, silyloxyalkyl, aralkyl, cycloalkylcarbonyloxyalkyl, arylalkenyl, arylcarbonyloxyalkyl, arylcarbonyl, aryloxycarbonyl, aralkoxycarbonyl, arylthioalkyl, aralkanoyl, aroyl, carboxyl, formyl, alkenylcarbonyl, alkynylcarbonyl, Het¹oxyalkyl, Het¹alkoxycarbonyl, Het¹oxycarbonyl, Het¹aryloxycarbonyl. Het¹alkyloxyalkyl. Het¹arylthioalkyl, Het¹aryloxyalkyl, Het oxyalkylcarbonyl, Het¹alkyloxyalkylcarbonyl, Het¹aralkoxycarbonyl, Het¹aryloxyalkylcarbonyl, Het¹carbonyloxyalkyl, Het¹alkylcarbonyloxyalkyl, Het²oxycarbonyl, Het²alkyloxyalkyl, Het²oxyalkyl, Het¹aralkylcarbonyloxyalkyl, Het²aralkoxycarbonyl, Het²aryloxycarbonyl, Het²aryloxyalkyl, Het²alkoxycarbonyl, Het²arylthioalkyl, Het²oxyalkylcarbonyl, Het²alkyloxyalkylcarbonyl, Het²aryloxyalkylcarbonyl, Het²carbonyloxyalkyl, Het²alkylcarbonyloxyalkyl, Het²aralkylcarbonyloxyalkyl, CR⁸=NR⁷, $CR^6=N(OR^7)$,

with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹ alkyl, Het¹ aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R² and R³ are independently selected from the group comprising hydroxyl, alkyloxy, alkyloxyalkyloxy, cycloalkyloxy cycloalkyloxy, aralkyloxy, aryloxyalkyloxy,





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silyloxy, alkylcarbonyloxy, arylcarbonyloxy, cycloalkylcarbonyloxy, aryloxycarbonylalkyloxy, formyloxy, Het¹alkyloxy, Het¹oxy, Het¹oxy, Het¹oxyalkyloxy, Het¹aryloxy, Het¹aralkyloxy, Het¹carbonyloxy, Het¹alkanoyloxy, Het¹aralkanoyloxy, Het¹aryloxyalkyloxy, Het²oxyalkyloxy, Het²aralkyloxy, Het²cycloalkyloxy, Het²alkanoyloxy, Het²aralkanoyloxy, Het²aryloxy, Het²aryloxy,

wherein R¹ R² and R³ are optionally substituted by one or more substituents independently selected from the group indicated in claim 1; and wherein R⁴ is oxo and R⁵ is hydrogen or alkyl.

5. A compound according to claim 1, 2 or 4,

wherein R¹ is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxyalkyl, alkylthioalkyl, alkanoyl, cycloalkylalkyl, cycloalkylalkyl, cycloalkylalkanoyl, cycloalkylalkyl, aralkyl, arylalkenyl, arylcarbonyl, arylthioalkyl, aralkanoyl, aroyl, carboxyl, formyl, alkenylcarbonyl, alkynylcarbonyl, Het¹axylakyl, Het¹aryloxyalkyl, Het¹arylthioalkyl, Het¹arylthioalkyl, Het¹aryloxyalkylcarbonyl, Het¹aryloxyalkylcarbonyl, Het²aryloxyalkylcarbonyl, Het²aryloxyalkyl, Het²aryloxyalkyl, Het²aryloxyalkyl, Het²aryloxyalkyl, Het²aryloxyalkyl, Het²aryloxyalkyl, CR⁵=N(OR²),

with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹ Het¹ alkyl, Het¹ aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R² and R³ are independently selected from the group comprising hydroxyl, alkylcarbonyloxy, arylcarbonyloxy, cycloalkylcarbonyloxy, formyloxy, Het¹carbonyloxy, Het²aralkanoyloxy, Het²aralkanoyloxy, Het²aralkanoyloxy,

wherein R¹ R² and R³ are optionally substituted by one or more substituents independently selected from the group indicated in claim 1; and

wherein R⁴ is oxo and R⁵ is hydrogen or alkyl.

6. A compound according to any of claims 1, 2, 4 to 5, wherein R¹ is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxyalkyl, alkylthioalkyl, cycloalkylalkyl, cycloalkylthioalkyl, silyloxyalkyl, aralkyl, arylalkenyl, arylthioalkyl, carboxyl, formyl, Het¹oxyalkyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het¹arylthioalkyl, Het²oxyalkyl, Het²aryloxyalkyl, Het²arylthioalkyl, optionally substituted by one or more





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substituents independently selected from the group indicated in claim 1; wherein R² and R³ are hydroxyl and wherein R⁴ is oxo and R⁵ is hydrogen.

- 7. A compound according to any of claims 1, 2, 4 to 6, wherein R¹ is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxyalkyl, cycloalkylalkyl, silyloxyalkyl, aralkyl, arylalkenyl, carboxyl, formyl, Het¹oxyalkyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het²oxyalkyl, Het²aryloxyalkyl, optionally substituted by one or more substituents independently selected from the group indicated in claim 1; wherein R² and R³ are hydroxyl, R⁴ is oxo₁and R⁵ is hydrogen.
- 8. A compound according to any of claims 1, 2, 4 to 7, wherein R¹ is selected from the group comprising alkyl, carboxyl, formyl; wherein R² and R³ are hydroxyl, and wherein R⁴ is oxo and R⁵ is hydrogen.
- 9. A compound according to claim 8, wherein R¹ is formyl, R² and R³ are hydroxyl R⁴ is oxo and R⁵ is hydrogen.
 - 10. A compound according to claim 1 or 3,

wherein R¹ is selected from the group comprising hydrogen, alkyl, alkenyl, alkynyl, alkyloxyalkyl, hydroxyalkyl, alkylthioalkyl, alkanoyl, cycloalkylalkyl, cycloalkylcarbonyl, cycloalkylalkanoyl, cycloalkylthioalkyl, silyloxyalkyl, aralkyl, arylalkenyl, arylcarbonyl, arylthioalkyl, aralkanoyl, aroyl, carboxyl, formyl, alkenylcarbonyl, alkynylcarbonyl, Het¹aryloxyalkyl, Het¹aryloxyalkyl, Het¹aryloxyalkyl, Het¹aryloxyalkyl, Het¹aryloxyalkyl, Het²aryloxyalkyl, CR6=NR7, CR6=N(OR7),

with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹ alkyl, Het¹ alkyl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R² and R³ are independently selected from the group comprising hydroxyl, alkylcarbonyloxy, arylcarbonyloxy, cycloalkylcarbonyloxy, formyloxy, Het¹carbonyloxy, Het¹alkanoyloxy, Het²aralkanoyloxy, Het²aralkanoyloxy,

wherein R¹ R² and R³ are optionally substituted by one or more substituents independently selected from the group indicated in claim 1; and



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wherein R⁴ is oxo, hydroxyalkyl, alkyl, alkenyl, arylcarbonylaryl, alkylcarbonylalkyl and R⁵ is hydrogen or alkyl.

- 11. A compound according to any of claims 1, 3 or 10, wherein R¹ is hydroxyalkyl, R² and R³ are hydroxyl, R⁴ is oxo and R⁵ is hydrogen.
- 12. A compound according to any of claims 1, 3 or 10, wherein R¹ is selected from the group comprising hydrogen, alkyl, alkenyl, alkynyl, hydroxyalkyl, alkyloxyalkyl, alkylthioalkyl, cycloalkylalkyl, cycloalkylthioalkyl, silyloxyalkyl, aralkyl, arylalkenyl, arylthioalkyl, carboxyl, formyl, Het¹oxyalkyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het¹arylthioalkyl, Het²oxyalkyl, Het²arylthioalkyl, optionally substituted by one or more substituents independently selected from the group indicated in claim 1; wherein R² and R³ are hydroxyl and wherein R⁴ is hydroxyalkyl, arylcarbonylalkyl, alkylcarbonylalkyl and R⁵ is hydrogen.
- 13. A compound according to any of claims 1, 3, 10 or 12, wherein R¹ is selected from the group comprising hydrogen, alkyl, alkenyl, alkynyl, hydroxyalkyl, alkyloxyalkyl, cycloalkylalkyl, silyloxyalkyl, aralkyl, arylalkenyl, carboxyl, formyl, Het¹oxyalkyl, Het¹aryloxyalkyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het²alkyloxyalkyl, Het²aryloxyalkyl, optionally substituted by one or more substituents independently selected from the group indicated in claim 1; wherein R² and R³ are hydroxyl, R⁴ is hydroxyalkyl, arylcarbonylalkyl, alkylcarbonylalkyl and R⁵ is hydrogen.
- 14. A compound according to any of claims 1, 3, 10, 12 or 13, wherein R¹ is selected from the group comprising alkyl, hydroxyalkyl, carboxyl, formyl; wherein R² and R³ are hydroxyl, and wherein R⁴ is arylcarbonylalkyl and R⁵ is hydrogen.
 - 15. A compound according to claim 14, wherein R¹ is hydroxyalkyl, R² and R³ are hydroxyl, R⁴ is arylcarbonylalkyl and R⁵ is hydrogen.
 - 16. A compound according to claim 15, wherein R¹ is hydroxymethylene, R² and R³ are hydroxyl, R⁴ is phenylcarbonylmethylene and R⁵ is hydrogen.



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17. A compound having the formula la or a pharmaceutically acceptable salt or ester thereof,

formula la

$$R_4$$
 R_5
 R_2
 R_1
 R_3
 R_4
 R_4
 R_5
 R_4
 R_5
 R_4
 R_4
 R_4
 R_5

wherein R1 is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxyalkyl, alkanoyl, cycloalkylcarbonyl, cycloalkylalkyl, alkylthioalkyl, alkyloxycarbonyl, cycloalkylaikoxycarbonyl, cycloalkylthioalkyl, alkylcarbonyloxyalkyl, cycloalkylalkanoyl, aralkyl, silyloxyalkyl, arylcarbonyloxyalkyl, cycloalkylcarbonyloxyalkyl, arylalkenyl, arylcarbonyl, aryloxycarbonyl, aralkoxycarbonyl, arylthioalkyl, aralkanoyl, aroyl, silyloxyalkyl, alkenylcarbonyl, Het¹oxyalkyl, Het¹alkoxycarbonyl, alkynyicarbonyl, carboxyl, Het¹oxycarbonyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het¹arylthioalkyl, Het¹aryloxycarbonyl, Het¹alkyloxyalkylcarbonyl, Het¹oxyalkylcarbonyl, Het¹aralkoxycarbonyl, Het¹alkylcarbonyloxyalkyl, Het¹carbonyloxyalkyl, Het¹aryloxyalkylcarbonyl, Het²alkyloxyalkyl, Het²oxyalkyl, Het²oxycarbonyl, Het¹aralkylcarbonyloxyalkyl, Het²aryloxycarbonyl, Het²alkoxycarbonyl, Het²aralkoxycarbonyl, Het²aryloxyalkyl, Het²arylthioalkyl, Het²oxyalkylcarbonyl, Het²alkyloxyalkylcarbonyl, Het²aryloxyalkylcarbonyl, Het²carbonyloxyalkyl, Het²alkylcarbonyloxyalkyl, Het²aralkylcarbonyloxyalkyl,CR⁶=NR⁷, $CR^6=N(OR^7)$,

with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹ Het¹ alkyl, Het¹ aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R² and R³ have the same definition as in claim 1;

wherein R¹ R² and R³ are optionally substituted by one or more substituents independently selected from the group as indicated in claim 1, and

wherein R⁴ and R⁵ are hydrogen or alkyl.

18. A compound according to claim 17,







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wherein R1 is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxyalkyl, alkylthioalkyl, alkanoyl, cycloalkylalkyl, cycloalkylcarbonyl, cycloalkylalkanoyl, cycloalkylthioalkyl, silyloxyalkyl, aralkyl, arylalkenyl, arylcarbonyl, arylthioalkyl, aralkanoyl, aroyl, silyloxyalkyl, carboxyl, alkenylcarbonyl, alkynylcarbonyl, Het¹oxyalkyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het¹arylthioalkyl, Het¹oxyalkylcarbonyl, Het¹alkyloxyalkylcarbonyl, Het¹aryloxyalkylcarbonyl, Het²oxyalkyl, Het²alkyloxyalkyl, Het²aryloxyalkyl, Het²arylthioalkyl, Het²oxyalkylcarbonyl, Het²alkyloxyalkylcarbonyl, Het²aryloxyalkylcarbonyl, CR⁶=N(OR⁷), with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹, Het¹alkyl, Het¹aryl, alkenyl, alkynyl, aminoalkyl, alkylcarbonylamino, aminoaryl, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R² and R³ have the same definition as in claim 1;

wherein R¹ R² and R³ are optionally substituted by one or more substituents independently selected from the group as indicated in claims 1, and

wherein R⁴ and R⁵ are hydrogen or alkyl.

- A compound according to claim 17 or 18, wherein R1 is selected from the group 19. comprising alkyl, alkenyl, alkyloxyalkyl, alkylthioalkyl, alkynyl, cycloalkylalkyl, cycloalkylthioalkyl, silyloxyalkyl, aralkyl, arylalkenyl, arylthioalkyl, silyloxyalkyl, carboxyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het'oxvalkyl. Het¹arvlthioalkvl. Het²oxvalkvl. Het²alkyloxyalkyl, Het²aryloxyalkyl, Het²arylthioalkyl, optionally substituted by one or more substituents independently selected from the group indicated in claim 1; wherein R² and R³ are hydroxyl and wherein R⁴ and R⁵ are hydrogen or alkyl.
- 25 20. A compound according to any of claims 17 to 19, wherein R¹ is selected from the group comprising alkyl, alkenyl, alkynyl, alkyloxyalkyl, cycloalkylalkyl, silyloxyalkyl, aralkyl, arylalkenyl, carboxyl, Het¹oxyalkyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het²oxyalkyl, Het²alkyloxyalkyl, Het²aryloxyalkyl, optionally substituted by one or more substituents independently selected from the group indicated in claim 1; wherein R² and R³ are hydroxyl and wherein R⁴ and R⁵ are hydrogen.
 - 21. A compound having the formula lb or a pharmaceutically acceptable salt or ester thereof,











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formula lb

$$R_4$$
 R_5
 R_2
 R_1
 R_3
 R_4
 R_4
 R_5
 R_4
 R_5
 R_1
 R_3

wherein R1 is selected from the group comprising alkenyl, alkynyl, alkyloxyalkyl, alkylthioalkyl, alkyloxycarbonyl, alkanoyl, cycloalkylalkyl, cycloalkylcarbonyl, cycloalkylalkanoyl, cycloalkylalkoxycarbonyl, cycloalkylthioalkyl, alkylcarbonyloxyalkyl, arylcarbonyloxyalkyl, cycloalkylcarbonyloxyalkyl, silyloxyalkyl, aralkyl, arylalkenyl, arylcarbonyl, aryloxycarbonyl, aralkoxycarbonyl, arylthioalkyl, aralkanoyl, aroyl, silyloxyalkyl, Het¹alkoxycarbonyl, Het¹oxyalkyl, carboxyl, alkenylcarbonyl, alkynylcarbonyl, Het¹oxycarbonyl, Het¹aryloxyalkyl, Het¹alkyloxyalkyl, Het¹arylthioalkyl, Het¹aryloxycarbonyl, Het¹oxyalkylcarbonyl, Het¹aralkoxycarbonyl, Het¹alkyloxyalkylcarbonyl, Het¹aryloxyalkylcarbonyl, Het¹carbonyloxyalkyl, Het¹alkylcarbonyloxyalkyl, Het¹aralkylcarbonyloxyalkyl, Het²oxyalkyl, Het²alkyloxyalkyl, Het²oxycarbonyl, Het²alkoxycarbonyl, Het²aralkoxycarbonyl, Het²aryloxycarbonyl, Het²aryloxyalkyl, Het²arylthioalkyl, Het²oxyalkylcarbonyl, Het²alkyloxyalkylcarbonyl, Het²aryloxyalkylcarbonyl, Het²carbonyloxyalkyl, Het²alkylcarbonyloxyalkyl, Het²aralkylcarbonyloxyalkyl, CR⁶=NR⁷. $CR^6=N(OR^7)$,

with R⁶ and R⁷ being independently selected from the group comprising hydrogen, hydroxyl, alkyl, aryl, Het¹, Het¹alkyl, Het¹aryl, alkenyl, alkynyl, aminoalkyl, aminoaryl, alkylcarbonylamino, arylcarbonylamino, alkylthiocarbonylamino and arylthiocarbonylamino;

wherein R¹ Is optionally substituted by one or more substituents independently selected from the group as indicated in claim 1, and

wherein R² and R³ are hydroxyl and wherein R⁴ is replaced by a double bond between the N atom and the C carbon atom of the N-containing heterocyclic ring of formula I; and wherein R⁵ is hydrogen.

22. A compound according to claim 21, wherein R¹ is selected from the group comprising alkenyl, alkyloxyalkyl, cycloalkylalkyl, silyloxyalkyl, aralkyl, arylalkenyl, carboxyl,

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Het oxyalkyl, Het aryloxyalkyl, Het alkyloxyalkyl, Het alkyloxyalkyl, Het alkyloxyalkyl, Het aryloxyalkyl, optionally substituted by one or more substituents independently selected from the group indicated in claim 1; wherein R² and R³ are hydroxyl and wherein R⁴ and R⁵ are hydrogen.

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23. A compound according to claim 22, wherein R¹ has the same definition as in claim 20, wherein R² and R³ are hydroxyl; wherein R⁴ is replaced by a double bond between the N atom and the C carbon atom of the N-containing heterocyclic ring of formula I; and wherein R⁵ is hydrogen.

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- 24. Compound of formula I, wherein R¹ is hydroxyalkyl, wherein R² and R³ are hydroxyl; wherein R⁴ is replaced by a double bond between the N atom and the C carbon atom of the N-containing heterocyclic ring of formula I; and wherein R⁵ is hydrogen.
- 25. Compound of formula I or a pharmaceutically acceptable salt or ester thereof, wherein R¹, R², R³, R⁴ and R⁵ are selected as in Table A.
 - 26. A pharmaceutical composition comprising a pharmaceutically acceptable excipient and a therapeutically effective amount of a compound according to any of claims 1-25.

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- 27. A pharmaceutical composition comprising a pharmaceutically acceptable excipient and a therapeutically effective amount of a compound according to claim 9.
- 28. A pharmaceutical composition comprising a pharmaceutically acceptable excipient and a therapeutically effective amount of a compound according to claim 11.
 - 29. A compound according to any of claims 1 to 25 for use as a medicament.
- 30. Use of a compound according to any of claims 1 to 25 for the preparation of a medicament for treating cancer.
 - 31. Use of a compound according to any of claims 1 to 25 in the treatment of cancer.

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32. Method of treating cancer comprising administrating to an individual in need of such treatment a pharmaceutical composition according to any of claims 26 to 28.



